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APPLICATION NO.		FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO	
10/049,822	•	04/01/2002	Shigeo Ohta	2002_0256A	6325	
513	7590	07/19/2006		EXAMINER		
	•	ND & PONACK, I	YAEN, CHRISTOPHER H			
2033 K STREET N. W. SUITE 800			ART UNIT	PAPER NUMBER		
WASHINGTON, DC 20006-1021				1643		
				DATE MAILED: 07/19/2006		

Please find below and/or attached an Office communication concerning this application or proceeding.

		Application No.	Applicant(s)				
		10/049,822	OHTA ET AL.				
	Office Action Summary	Examiner	Art Unit				
		Christopher H. Yaen	1643				
- The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply							
WHIC - Exter after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR REPLY CHEVER IS LONGER, FROM THE MAILING DANS of time may be available under the provisions of 37 CFR 1.13 SIX (6) MONTHS from the mailing date of this communication. Operiod for reply is specified above, the maximum statutory period we to reply within the set or extended period for reply will, by statute, reply received by the Office later than three months after the mailing ed patent term adjustment. See 37 CFR 1.704(b).	ATE OF THIS COMMUNICATION 36(a). In no event, however, may a reply be tim rill apply and will expire SIX (6) MONTHS from the cause the application to become ABANDONEI	I. sely filed the mailing date of this communication. O (35 U.S.C. § 133).				
Status							
1)⊠	Responsive to communication(s) filed on 21 Ap	<u>oril 2006</u> .					
2a) <u></u> □	This action is FINAL . 2b)⊠ This action is non-final.						
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is						
closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.							
Dispositi	on of Claims						
5)□ 6)⊠ 7)□	Claim(s) <u>1-25</u> is/are pending in the application. 4a) Of the above claim(s) <u>6-8,18-20 and 25</u> is/a Claim(s) is/are allowed. Claim(s) <u>1-5,9-17 and 21-24</u> is/are rejected. Claim(s) is/are objected to. Claim(s) are subject to restriction and/or						
Application Papers							
10)□	The specification is objected to by the Examiner The drawing(s) filed on is/are: a) access applicant may not request that any objection to the correction of the correct	epted or b) objected to by the Edrawing(s) be held in abeyance. See on is required if the drawing(s) is objected to be a second to be a secon	e37 CFR 1.85(a). ected to. See 37 CFR 1.121(d).				
Priority u	ınder 35 U.S.C. § 119						
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 							
Attachment	(s)						
	e of References Cited (PTO-892)	4) Interview Summary (PTO-413)				
3) 🔲 Infom	e of Draftsperson's Patent Drawing Review (PTO-948) nation Disclosure Statement(s) (PTO-1449 or PTO/SB/08) r No(s)/Mail Date	Paper No(s)/Mail Dai 5) Notice of Informal Pa 6) Other:	te atent Application (PTO-152)				

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DETAILED ACTION

Re: OHTA ET AL

1. The amendment filed 4/26/2006 is acknowledged and entered into the record.

2. Claims 1-25 are pending, claims 6-8,18-20, and 25 are withdrawn as being

drawn to a non-elected invention.

3. Claims 1-5,9-17, and 21-24 are examined on the merits.

4. The text of those sections of Title 35, U.S. Code not included in this action can

be found in a prior Office action.

Claim Objections

5. Claims 1-5, 9-17, and 21-24 are objected to because of the following

informalities: the claims recite the terms "Tyr", "Phe", "Gln", "Asn", "Arg", and "Lys" to

represent amino acids tyrosine, phenylalanine, glutamine, asparagine, arginine, and

lysine. The first appearance of an abbreviation must be accompanied by the full name

of the term. Appropriate correction is required.

NEW REJECTIONS

Claim Rejections - 35 USC § 112, 1st paragraph

6. Claims 1-5 and 9-12 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the written description requirement. The claim(s) contains subject matter

which was not described in the specification in such a way as to reasonably convey to

one skilled in the relevant art that the inventor(s), at the time the application was filed,

had possession of the claimed invention. The written description in this case has only set forth a nucleic acid sequence comprising "the" nucleotide sequence of SEQ ID No: 1, except having substitutions wherein the nucleic acid sequences encode amino acid substitutions of Tyr 22 to Phe, Gln 26 to Asn, Arg 165 to Lys, and therefore the written description is not commensurate in scope to the claims that read on a peptide sequence which consists of or comprises "a" sequence of SEQ ID No: 1 with such changes. The following written description rejection is set forth herein.

The claims recite "a" nucleotide sequence of SEQ ID No: 1 as part of the

invention. This reads on a single amino acid found within the sequence of SEQ ID No:

1. However, there does not appear to be an adequate written description in the specification as-filed that is representative of the single amino acid sequences derived from SEQ ID No: 1, which is encompassed by the claimed peptide sequences. The Guidelines for the Examination of Patent Applications Under the 35 U.S.C. 112, ¶ 1 "Written Description" Requirement make clear that the written description requirement for a claimed genus may be satisfied through sufficient description of a representative number of species by actual reduction to practice, reduction to drawings, or by disclosure of relevant, identifying characteristics, i.e., structure or other physical and or chemical properties, by functional characteristics coupled with a known or disclosed correlation between function and structure, or by a combination of such identifying characteristics, sufficient to show the applicant was in possession of the genus (Federal Register, Vol. 66, No. 4, pages 1099-1111, Friday January 5, 2001, see especially page 1106 3rd column).

Applicant does not appear to have reduced to practice the broad genus of "a" nucleotide sequence of SEQ ID No: 1 derived from SEQ ID No: 1. Neither has Applicant provided a sufficient written description of any particular structure of "a" nucleotide sequence derived from SEQ ID No: 1. "[A] nucleotide sequence" encompasses any nucleotide, as small as 2 nucleotides, found within SEQ ID No: 1. Thus the genus of compounds encompassed by this phrase is extensive and the artisan would not be able to recognize that Applicant was in possession of the invention as now claimed.

Consequently, Applicant was not in possession of the instant claimed invention.

See Regents of the University of California v. Eli Lilly and Co. 119 F.3d 1559, 43

USPQ2d 1398 (Fed. Cir. 1997). Adequate written description of genetic material

"requires a precise definition, such as by structure, formula, chemical name, or physical properties,' not a mere wish or plan for obtaining the claimed chemical invention." Id. 43

USPQ2d at 1404 (quoting Fiers, 984 F.2d at 1171, 25 USPQ2d at 1606). The disclosure must allow one skilled in the art to visualize or recognize the identity of the subject matter of the claim. Id. 43 USPQ2d at 1406. A description of what the genetic material does, rather than of what it is, does not suffice. Id.

Applicant is directed to the Guidelines for the Examination of Patent Applications Under the 35 U.S.C. 112, ¶ 1 "Written Description" Requirement, Federal Register, Vol. 66, No. 4, pages 1099-1111, Friday January 5, 2001. Applicant is also invited to point to clear support or specific examples of the claimed invention in the specification asfiled.

It is noted that applicant may overcome this rejection by amending the claims to recite a sequence comprising "the" nucleotide sequence.

Claim Rejections - 35 USC § 112, 1st paragraph

7. Claims 1-5, 9-17, and 21-24 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the enablement requirement. The claim(s) contains subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to use the invention.

Factors to be considered in determining whether a disclosure meets the enablement requirement of 35 USC 112, first paragraph, have been described by the court in *In re Wands*, 8 USPQ2d 1400 (CA FC 1988). Wands states at page 1404,

"Factors to be considered in determining whether a disclosure would require undue experimentation have been summarized by the board in Ex parte Forman. They include (1) the quantity of experimentation necessary, (2) the amount of direction or guidance presented, (3) the presence or absence of working examples, (4) the nature of the invention, (5) the state of the prior art, (6) the relative skill of those in the art, (7) the predictability or unpredictability of the art, and (8) the breadth of the claims."

The nature of the invention

The claims are drawn to genetically engineered cDNAs, vectors comprising said cDNA, and host cells comprising said cDNA. The invention is in a class of invention which the CAFC has characterized as "the unpredictable arts such as chemistry and biology." *Mycogen Plant Sci., Inc. v. Monsanto Co.*, 243 F.3d 1316, 1330 (Fed. Cir. 2001).

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The breadth of the claims

The claims encompass products that are intended to be used for administration to induce cells into an apoptotic mechanism and therefore read on gene therapy.

The unpredictability of the art and the state of the prior art

At the time the application was filed, the art of administering any type of genetic expression vector to an individual so as to provide a tangible therapeutic benefit was poorly developed and unpredictable. The NIH ad hoc committee to assess the current status and promise of gene therapy reported in December 1995 that "clinical efficacy has not been definitively demonstrated at this time in any gene therapy protocol, despite anecdotal claims...," and that "significant problems remain in all basic aspects of gene therapy" (Orkin and Motulsky, p. 1). In a review article published in Scientific American in June 1997, Theodore Friedmann discusses the technical barriers which have so far prevented successful gene therapy, and states "So far, however, no approach has definitively improved the health of a single one of the more than 2,000 patients who have enrolled in gene therapy trials worldwide" (p. 96). In a review article published in Nature in September 1997, Inder Verma states "Although more than 200 clinical trials are currently underway worldwide, with hundreds of patients enrolled, there is still no single outcome that we can point to as a success story" (p. 239).

In an article published well after the effective filing date of the instant application, Rubanyi (2001) teaches that the problems described above remain unsolved at the time the instant application was filed. Rubanyi states, "[a]Ithough the theoretical advantages of [human gene therapy] are undisputable, so far [human gene therapy] has not

delivered the promised results: convincing clinical efficacy could not be demonstrated yet in most of the trials conducted so far ..." (page 113, paragraph 1). Among the technical hurdles that Rubanyi teaches remain to be overcome are problems with gene delivery vectors and improvement in gene expression control systems (see especially the section under "3. Technical hurdles to be overcome in the future", pp. 116-125).

The state of the art is such that no correlation exists between successful expression of a gene and a therapeutic result (Ross et al., p. 1789, column 1, paragraph 1).

Working examples

The working examples of the instant invention are drawn to the administration of a protein encoded by the instantly claimed invention or to a methods of making the instantly claimed invention.

Guidance in the specification

The specification fails to teach how the administration of the instant invention can be used as a means of altering the apoptotic mechanism or pathway as intended through gene therapeutic methods. Moreover, the specification provides little guidance for the use of the claimed invention for any method or purpose associated with gene therapy, given the generally unpredictable nature of the art pertaining to gene therapy.

Level of skill in the art

The level of skill in the art is deemed to be high.

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Conclusion

Thus given the broad claims in an art whose nature is identified as unpredictable, the unpredictability of that art, the large quantity of research required to define these unpredictable variables, the lack of guidance provided in the specification, the presence of a working example which does not address the issue of the efficacy of the control and the negative teachings in the prior art balanced only against the high skill level in the art, it is the position of the examiner that it would require undue experimentation for one of skill in the art to perform the method of the claim as broadly written.

Conclusion

No claim is allowed.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Christopher H. Yaen whose telephone number is 571-272-0838. The examiner can normally be reached on Monday-Friday 9-5.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Larry Helms, Ph.D. can be reached on 571-272-0832. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

Christopher Yaen Art Unit 1643 July 6, 2006 Chrustopher H. Yaen Primary examiner